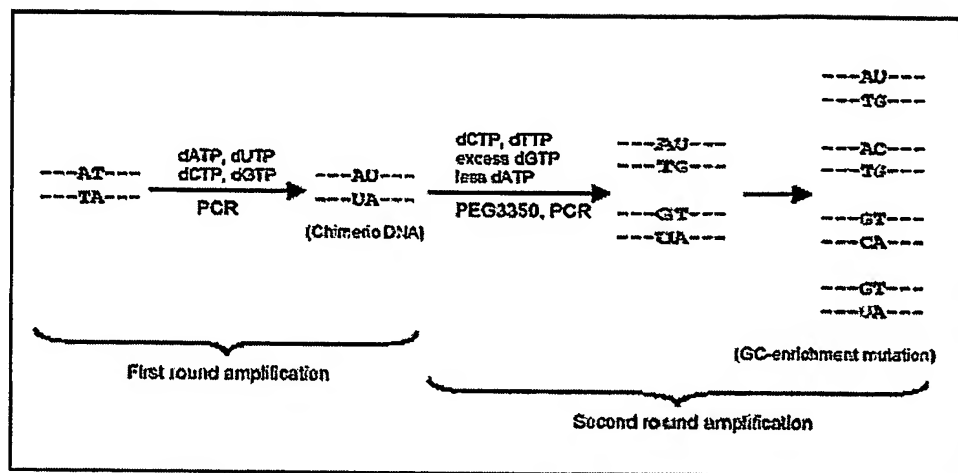


Figure 1



5

Figure 2

	34	38	41	103	105	107
AlbD	-R C I L F A R G L S G -	-	-	G V	G H S L G S V-	
Bhc	-R A V L L I H G F T G -	-	-	V G G L S L G G V-		
Sac	-R A V L L L I H G F T G -	-	-	V R G L S I L G G D-		
Lic	-R A V L L L I H G F T G -	-	-	I I G F S M G G V-		
Bsc	-K A V L L L I H G F T G -	-	-	A C G L S L G G V-		
Tme	-K G V L F I H G Y T G -	-	-	I I G F S M G G V-		
Tpc	-K A V L L I H G Y M G -	-	-	V G G L S M G G V-		
Ful	-Y P I I L Y K G L T G -	-	-	L V G H S Q G G Q-		
Bsc	-R P L V I L R G F T G -	-	-	I I A H S Q G G Q-		
Consensus	L x x x G x x G			G x S x G		
	(I)			(A)		
	(V)					

Figure 3

MDKSDLTETSRIKHGEAEFDVTLLQVKGATRCILFAAGLGGSPLRHLELLQ
TFARHGVSVVAPHFERLTSPVPTRAELLERCQRLARAQNEFCSGYASVTGV
5 GHSLGSVILLLNAGAIAMTSAGE SVVFAGDRMLHRLILLAPPADFFQAPSA
LAAVNVPVHIWAGEKDSLTPPSQACFLKQALEGYTQTYLCVMEEAGHFTFM
NTLPPQVTDSHPSREAFLLDLGENIARLVTD

Figure 4

10

ATGGACAAAAGTGATCTCACGGAAACGTCTCGGATCAAACATGGGAAGAG
GCGTTGACGTACCTTATTGCAGGTTAAGGGGGCGACGCGCTGTATCCTT
TTTGCCTGCGGGCTGGCGGCAGTCCGCTGCGCCATCTGAACCTCTCCAG
ACCTTGCCGCCATGGCGTTCCGTTGCGGCCACACTTGAACGGTTG
15 ACCTCACCGTGCCCACCAGAGCTGAATTACTGGAACGCTGCCAGCGGCTT
GCGCGGCTCAGAATGAATTGTAGCGTTATGCGTCGGTACCGGTGTT
GGCCACTCCCTGGTAGCGTGATTTATTGCTGAATGCCGGGCTATAGCG
ATGACAAGCGCAGGGGAATCGTTGTTTGCCTGGCGACCGGATGTTGCAT
CGACTTATTTACTGGCACCGCCGCCGATTTTCCAGGCTCCGTCTGCG
20 CTGGCAGCGGTGAACGTACCTGTTACATCTGGCAGGTGAAAAGGACAGC
CTGACGCCCGTCCCAGGCCTGCTTCTTAAACAGGCAGTGGAGGGTTAC
ACGCAGACTTATCTGTGTGATGGAAGAGGCCGGCATTTACCTTCATG
AATACCTTGCCCTCCGCAGGTAACCGATTCACATCCGTGCGGGAGGCCTT
CTTTAGATTGGCGAAAACATAGCCCAGGTGGTACTGAT

25